

Development of ferromagnetism in the doped topological insulator Bi_{2-x}Mn_xTe₃

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The development of ferromagnetism in Mn-doped Bi_2Te_3 is characterized through measurements on a series of single crystals with different Mn content. Scanning tunneling microscopy analysis shows that the Mn substitutes on the Bi sites, forming compounds of the type $Bi_{2-x}Mn_xTe_3$, and that the Mn substitutions are randomly distributed, not clustered. Mn doping first gives $Bi_{2-x}Mn_xTe_3$, and that the Mn substitutions are randomly distributed, not clustered. Mn doping first gives $Bi_{2-x}Mn_xTe_3$, a second-order ferromagnetic transition is observed, with $T_C \sim 9-12$ K. The easy axis of magnetization in the ferromagnetic phase is perpendicular to the Bi_2Te_3 basal plane. Thermoelectric power and Hall effect measurements show that the Mn-doped Bi_2Te_3 crystals are p-type. Angle-resolved photoemission spectroscopy measurements show that the topological surface states that are present in pristine Bi_2Te_3 are also present at 15 K in ferromagnetic Mn-doped $Bi_{2-x}Mn_xTe_3$ and that the dispersion relations of the surface states are changed in a subtle fashion.

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I. INTRODUCTION

Bi₂Te₃ has long been studied as the parent compound of a family of excellent ambient-temperature thermoelectric materials (see, e.g., Refs. 1 and 2). Recently, however, Bi₂Se₃ and Bi₂Te₃ have been of renewed interest because they are bulk topological insulators (see, e.g., Refs. 3-7), hosting Dirac-like conducting surface states. 8–10 The doping of topological insulators to decrease bulk carrier concentration¹¹ and induce superconductivity¹² has been performed in order to investigate various aspects of the surface states in these materials, including their potential applicability in novel electronic devices. Of particular interest in future work will be the effects of magnetic impurities and ferromagnetism on the topological surface states. Here we report the development of ferromagnetism near 12 K as a function of Mn doping in single crystals of Bi_{2-x}Mn_xTe₃. Our work builds on a previous report of ferromagnetism in Bi_{1.98}Mn_{0.02}Te₃ (Ref. 13) and other work on the dilute ferromagnetic V-VI semiconductors Bi_{2-x}Fe_xTe₃ (Ref. 14) (a very weak ferromagnet) and Sb_{2-x}V_xTe₃. 15 Ferromagnetism in Mn-doped III-V semiconductors has been widely studied [see, e.g., Ref. 16]; the behavior we observe in the current materials is analogous. We show quantitatively how the Bi_{2-x}Mn_xTe₃ system develops ferromagnetism as a function of doping, characterize the material at the nanoscale by scanning tunneling microscopy (STM), and present preliminary angle-resolved photoemission spectroscopy (ARPES) characterization of the effects of doping on the surface states.

II. EXPERIMENT

High purity elemental Bi (99.999%), Mn (99.99%), and Te (99.999%) were used for the $Bi_{2-x}Mn_xTe_3$ crystal growth, employing nominal x values of 0.005, 0.01, 0.02, 0.05, and 0.1. Because Mn reacts with quartz, a two-step melting method, described in Ref. 11, was necessary for successful modified Bridgeman crystal growth. The crystal growth for

Bi_{2-x}Mn_xTe₃ involved cooling from 950 to 550 °C over a period of 24 h and then annealing at 550 °C for 3 days; silver-colored single crystals were obtained. The crystals were confirmed to be single phase and identified as having the rhombohedral Bi₂Te₃ crystal structure by x-ray diffraction using a Bruker D8 diffractometer with Cu $K\alpha$ radiation and a graphite diffracted beam monochromator. X-ray diffraction patterns of the cleaved crystals oriented with basal plane normals bisecting the incident and diffracted beam directions showed only the (0 0 3), (0 0 6), and (0 0 15) peaks (hexagonal setting), indicating that the cleaved surface is oriented perpendicular to the hexagonal c axis. The c-axis parameter does not change substantially with Mn doping: we refine the values to be 30.488(2) Å for Bi₂Te₃ and 30.467(3) Å for the highest Mn-doped sample. Elemental analysis (Galbraith Laboratories) indicated that the Mn contents in the single crystals are very close to the nominal values, with the correspondences: nominal x=0.005, 0.01, 0.02, 0.05, and 0.1 are equal to the true x=0.005, 0.01, 0.02, 0.04, and 0.09 respectively. The analytically determined Mn concentrations are employed in this report. dc magnetization measurements were performed on a Quantum Design Physical Property Measurement System (PPMS). Temperaturedependent resistivity measurements were carried out in the PPMS using the standard four-point probe technique with silver paste cured at room temperature used for the contacts. Seebeck coefficient measurements were conducted using a homemade probe with an MMR Technologies SB100 Seebeck measurement system. Hall effect measurements were performed between 5 and 200 K in a home-built apparatus. In all cases, the electric and thermal currents were applied in the basal plane (hexagonal ab plane) of the crystals. The Bi_{2-r}Mn_rTe₃ crystal surfaces were characterized with a homemade cryogenic scanning tunneling microscope (STM) at 4.2 K under ultrahigh vacuum conditions. ARPES data were measured on in situ cleaved crystal surfaces at 15 K, in a vacuum maintained below 6×10^{-11} torr, at the Advanced Light Source, beam lines 10 and 12.

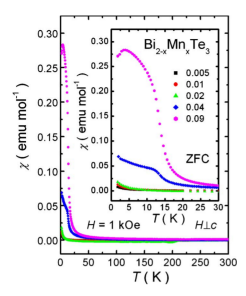


FIG. 1. (Color online) Zero-field cooled (ZFC) temperature-dependent dc magnetic susceptibility χ measured at 1 kOe applied magnetic field for the $\text{Bi}_{2-x}\text{Mn}_x\text{Te}_3$ crystals. The magnetic susceptibility in the region of 0 to 30 K is shown in the inset.

III. RESULTS AND DISCUSSION

The temperature-dependent magnetic susceptibilities, χ =M/H, measured in an applied field of 1 kOe [measured magnetizations (M) are linearly dependent on applied magnetic field (H) up to 1 kOe are shown in Fig. 1 for the $Bi_{2-x}Mn_xTe_3$ crystals for x=0, 0.005, 0.01, 0.02, 0.04, and0.09, measured with H perpendicular to the c axis. The susceptibility is negative at high temperatures for x < 0.04, an indication of the weakness of the paramagnetic contribution from the Mn, which becomes significant only at low temperatures. There is no indication of a ferromagnetic transition in the susceptibility measurements for x < 0.04. The system becomes ferromagnetic when the Mn concentration is increased to x=0.04, evidenced in these data by a relatively large susceptibility at low temperatures. The inset of Fig. 1 emphasizes the low temperature susceptibilities for the series; for the crystals with x=0.04 and 0.09 in $Bi_{2-x}Mn_xTe_3$, paramagnetic behavior is seen for the full temperature range between 300 K and T_C .

The susceptibility χ can be fit to the Curie-Weiss law, χ - χ_0 = $C/(T-\theta)$, where χ_0 is the temperature-independent term, C is the Curie constant, and θ is the Weiss temperature. Figure 2 shows the low-temperature inverse susceptibility plots, $1/(\chi-\chi_0)$ vs. T, for all samples. The inverse susceptibilities for x=0.005, 0.01, and 0.02 show straight line behavior, as expected for the presence of weak local moments due to the Mn dopants. For the lowest x, a very small negative $\theta(-0.7 \text{ K})$ is observed, which becomes slightly more positive with increasing Mn content. In contrast, data for crystals of composition x=0.04 and 0.09 follow the Curie-Weiss law with positive θ s of 11 and 13 K, respectively. Figure 3 shows the temperature dependence of the inverse susceptibility for the Bi_{2-x}Mn_xTe₃ crystals normalized on the basis of moles of Mn for x = 0.005, 0.01, 0.02, 0.04, and 0.09. From these data, the effective moment per Mn, p_{eff} can be obtained from

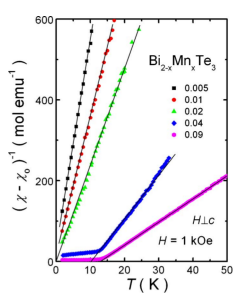


FIG. 2. (Color online) Temperature dependence of the inverse susceptibility for x=0.005, 0.01, 0.02, 0.04, and 0.09 of $Bi_{2-x}Mn_xTe_3$.

 p_{eff} =(7.99C)^{1/2}. As the Mn doping initially increases, the effective moment per mol-Mn decreases, but for most of the range, x=0.02–0.09, p_{eff} per Mn does not change much, as seen in the inset of Fig. 3, giving the magnitude of p_{eff} $\sim 4\mu_B$ per mol-Mn.

To better determine the temperatures of the ferromagnetic transitions, we apply the criterion employed for disordered systems 17,18 and present Arrott plots in Figs. 4 and 5 for Bi_{1.96}Mn_{0.04}Te₃ and Bi_{1.91}Mn_{0.09}Te₃. In such plots, H/M versus M^2 should go as $H/M = a'(T-T_C) + b'M^2 + c'M^4$ at values of H above those of the magnetic hysteresis. At T_C , the intercept changes sign. For second order transitions, b' is positive to the second order transitions, b' is positive transitions.

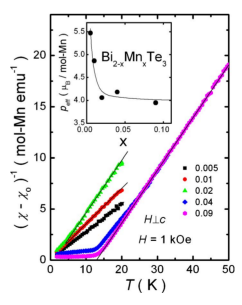


FIG. 3. (Color online) Temperature dependence of the inverse susceptibility per mole of Mn for x=0.005, 0.01, 0.02, 0.04, and 0.09 of $Bi_{2-x}Mn_xTe_3$. The effective magnetic moment per mole of Mn is shown in the inset.

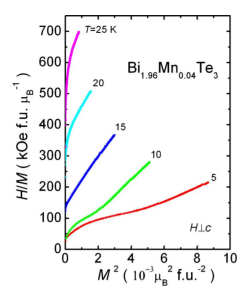


FIG. 4. (Color online) Arrott plots at various temperatures near T_C for Bi_{1.96}Mn_{0.04}Te₃. The ferromagnetic transition is of second order at $T_C \sim 9~$ K.

tive. Before each run, the samples were heated above their T_C and cooled to the measuring temperature under zero field to ensure perfect demagnetization. The data in Fig. 4 indicate that the ferromagnetic transition in $\mathrm{Bi}_{1.96}\mathrm{Mn}_{0.04}\mathrm{Te}_3$ is second order as expected, with T_C determined as ~ 9 K. The Arrott plot for $\mathrm{Bi}_{1.91}\mathrm{Mn}_{0.09}\mathrm{Te}_3$, Fig. 5, similarly shows a second-order ferromagnetic transition with a T_C of approximately 12 K.

Figure 6 shows the field-dependent magnetization curves at 1.8 K for a $\text{Bi}_{1.91}\text{Mn}_{0.09}\text{Te}_3$ crystal, where the applied field is parallel to the c axis, $H \parallel c$, and perpendicular to the c axis, $H \perp c$. The magnetization for $H \parallel c$ has a very steep initial slope, whereas the magnetization for $H \perp c$ increases much more slowly. This indicates that the c axis is the easy axis of magnetization, with spontaneously ordered magnetic mo-

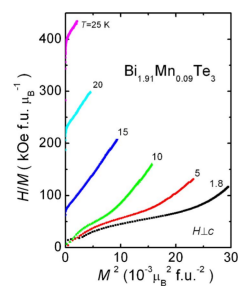


FIG. 5. (Color online) Arrott plots for $\mathrm{Bi}_{1.91}\mathrm{Mn}_{0.09}\mathrm{Te}_3$ show the second-order ferromagnetic transition with a T_C of about 12 K.

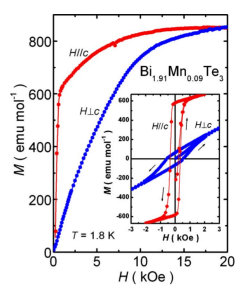


FIG. 6. (Color online) Magnetic-field-dependent magnetization of a $\text{Bi}_{1.91}\text{Mn}_{0.09}\text{Te}_3$ crystal under the applied field parallel to the c axis, $H \parallel c$, and perpendicular to the c axis, $H \perp c$, at $T = 1.8\,$ K. Inset shows the hysteresis MH loops of this $\text{Bi}_{1.91}\text{Mn}_{0.09}\text{Te}_3$ crystal for both field directions.

ments in the ferromagnetic phase aligned perpendicular to the Bi₂Te₃ basal plane. This moment direction is the same as is seen in V-doped Sb₂Te₃. ¹⁵ A narrow hysteresis loop is seen for $H \parallel c$, as shown in the inset of Fig. 6. The small coercivity, $H_C \sim 350$ Oe, indicates that the sample is a soft ferromagnet. H_C for our crystal of $Bi_{1.91}Mn_{0.09}Te_3$ is much smaller than is seen for V-doped Sb₂Te₃, where H_C is \sim 12 kOe at 2 K.¹⁵ The saturated magnetic moment, reached by about 12 kOe for $H \parallel c$ and 20 kOe for $H \perp c$ at 1.8 K, is 1.5 μ_B per mol Mn, which is much smaller than p_{eff} measured above T_C . This large decrease follows the pattern in itinerant ferromagnets with strongly suppressed T_C (as here). The likely reason is the very weak ferromagnetic exchange energy. In the ordered state, the suppressed exchange translates to a magnetization that is sharply reduced from that obtained by fully aligning the local moment of each magnetic ion. In MnSi, for example, ¹⁹ which becomes a spiral magnet at T_C =30 K, the high-field saturated magnetization at $T \ll T_C$ gives $0.5 \mu_B$ per Mn even though the Curie-Weiss susceptibility measures a local moment of $3-4\mu_B$ above T_C .

The temperature dependence of the basal plane resistivities for $\mathrm{Bi}_{2-x}\mathrm{Mn}_x\mathrm{Te}_3$ crystals for x=0.005, 0.02, and 0.09 are shown in Fig. 7. As Mn-doping increases, the resistivity of $\mathrm{Bi}_{2-x}\mathrm{Mn}_x\mathrm{Te}_3$ increases, though the metallic behavior typically observed in heavily doped small band gap semiconductors is maintained. The increased resistivity at low temperatures in the more heavily doped samples is due to the presence of a larger residual resistivity, indicative of strong carrier scattering from the doped Mn. There is a sharp drop in resistivity for $\mathrm{Bi}_{1.91}\mathrm{Mn}_{0.09}\mathrm{Te}_3$ at ~ 13 K that accompanies the ferromagnetic transition seen in the susceptibility measurements. The resistivity anomaly is pushed to higher temperature and substantially broadened under a 10 kOe magnetic field applied parallel to the c axis, shown in the inset of Fig. 7. Broad drops in the resistivities for x=0.02 and 0.09

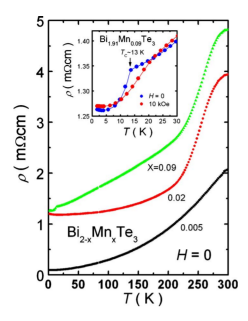


FIG. 7. (Color online) Temperature-dependent resistivities for x=0.005, 0.02, and 0.09 of $\mathrm{Bi}_{2-x}\mathrm{Mn}_x\mathrm{Te}_3$. Inset shows the resistivity of $\mathrm{Bi}_{1.91}\mathrm{Mn}_{0.09}\mathrm{Te}_3$ crystal in 0 and 10 kOe applied magnetic fields.

crystals are observed at temperatures around 250 K. The origin of these resistivity drops is unknown; further investigation is suggested. From the high field (14 T) values of the Hall resistivity ρ_{vx} on crystals with x=0.04 and 0.09, we find that the carriers are holelike with Hall density $p=H/e\rho_{vx}$ =5×10¹⁸ and 7×10¹⁹ cm⁻³, respectively (Fig. 8). In the x =0.04 sample, weak quantum oscillations resolved in the derivative $d\rho_{vr}/dH$ can be observed as depicted in Fig. 9. The period of the oscillations, $\Delta(1/H)=0.046~\mathrm{T}^{-1}$, corresponds to a small FS pocket with Fermi wave vector k_F =0.047 Å⁻¹ and carrier density 3.5×10^{18} cm⁻³ (assuming a spherical Fermi surface), a value slightly smaller than the measured Hall density. Figure 10 shows the thermoelectric power as a function of temperature for the Bi_{2-x}Mn_xTe₃ crystals. Bi_{2-x}Mn_xTe₃ is p-type in the whole range of temperatures for Mn substitutions up to x=0.09. Data in Fig. 10 show that Mn doping on the Bi site dramatically reduces the Seebeck coefficient, from 280 to 80 μ V K⁻¹ at room temperature. As temperature decreases, the Seebeck coefficients for Mn-doped Bi₂Te₃ monotonically decrease.

In order to directly visualize the effects of Mn doping in Bi₂Te₃ at the atomic scale, we have characterized the Bi_{2-x}Mn_xTe₃ (001) surface with a cryogenic STM. Due to the weak van der Waals bonding between the adjacent Te layers, the uppermost layer exposed after in situ cleavage is a triangular lattice of Te atoms. A typical STM topographic image of the surface of Bi_{1.91}Mn_{0.09}Te₃ is presented in Fig. 11(a). The black triangles are identified as substitutional Mn. Figures 11(b) and 11(c) show zoomed-in views over a triangular local density of states (LDOS) suppression associated with Mn dopants, in which the position of Te atoms on the top most layer are marked by blue (dark) circles. Below the surface Te layer, the Bi atoms form an interpenetrating triangular lattice [sites denoted by red (medium) circles] centered in the middle of the triangle formed by the Te atoms in the surface layer. The triangular suppression in the LDOS that

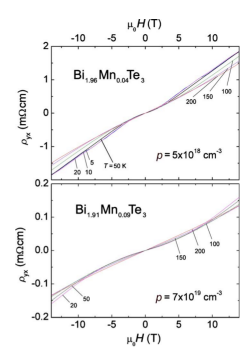


FIG. 8. (Color online) Hall resistivity ρ_{yx} vs field H in Bi_{2-x}Mn_xTe₃ with x=0.04 (upper panel) and x=0.09 (lower panel). ρ_{yx} is holelike and displays a slight change of slope in the weak-field limit. The hole density p= $H/e\rho_{yx}$ inferred at 8 T and 20 K equals 5×10^{18} cm⁻³ and 7×10^{19} cm⁻³ for x=0.04 and 0.09, respectively.

extends over several lattice sites is in the same lattice position expected for a Bi atom; therefore, we conclude that Mn substitutes primarily for Bi upon doping. In the topographies of the filled states, as shown in Fig. 11(c), Mn dopants are also centered on the Bi sites, forming a smaller region of suppression and a bright halo signifying enhanced LDOS. We attribute the formation of this bright region to local band bending due to negative charge that localizes about the Mn ion. This is as expected around substitutional acceptor atoms. In addition to the Mn dopants in the Bi layer, faint cloverlike patterns are visible in topographic images. These could be

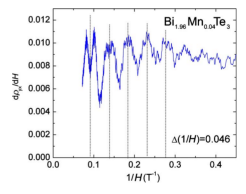


FIG. 9. (Color online) Quantum oscillations extracted from the derivative of the Hall resistivity $d\rho_{yx}/dH$ measured at 5 K in Bi_{2-x}Mn_xTe₃ with x=0.04. The period of the oscillations $\Delta(1/H)$ =0.046 T⁻¹ gives a (spherical) FS volume of wave vector k_F =0.047 Å⁻¹, corresponding to a carrier density, 3.5×10^{18} cm⁻³, slightly smaller than p= $H/e\rho_{yx}$.

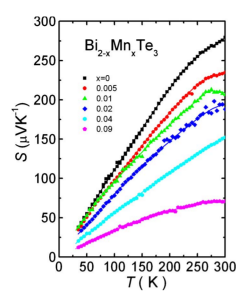


FIG. 10. (Color online) Temperature-dependent Seebeck coefficients for $Bi_{2-\nu}Mn_{\nu}Te_3$ crystals.

substitutional Mn atoms in deeper Bi layers, but their accurate identification requires further study. The number of contrast features associated with Mn atoms seen in the top most Bi layer appears to be significantly less than that expected from the measured Mn concentration; the reason for this is not known, but we speculate that the reduction of Mn concentration has resulted from the diffusion of the Mn dopants away from the surface layer after the cleavage. Such a diffusion process may be enhanced by the electric field which is created at the surface after cleaving. The implication of this scenario is that Mn may be intercalated between the Bi₂Te₃ layers in addition to substituting for Bi in the crystal lattice.

The STM topographic images can be used to study the spatial distribution of the Mn dopant atoms. Given the high concentration of Mn in ferromagnetic samples of Bi_{2-x}Mn_xTe₃, the formation of Mn clusters, if present, could indicate a completely different mechanism for the observed ferromagnetism. By locating the position of substitutional Mn, STM provides a direct visual method for examining the tendency to form clusters from the atomic to submicron scale. As discussed above, the substitutional Mn atoms in the closest Bi layer to the surface appear as triangular suppression of the LDOS in topographic images. After finding the position of the individual Mn dopants in one of such Bi layers, the correlation between Mn pairs was calculated for large fields of view (>1000 Å). The plot presented in Fig. 11(d) is the cumulative probability of random pair separation; i.e., for a given value of r, P(< r) is the probability of two randomly chosen Mn dopants being less than a distance r from one another. For homogeneously distributed dopants, this probability scales with the area (r^2) and the dopants fill the field of view, leaving no void area. The measured probability shows a power law behavior with a power very close to 2 for an extended range of distances (10 Å < r < 200 Å), indicating the absence of clustering in Bi_{2-x}Mn_xTe₃ for concentrations as high as x=0.09. In contrast to the single exponent observed here, the tendency toward cluster formation would result in different exponents at different length scales,

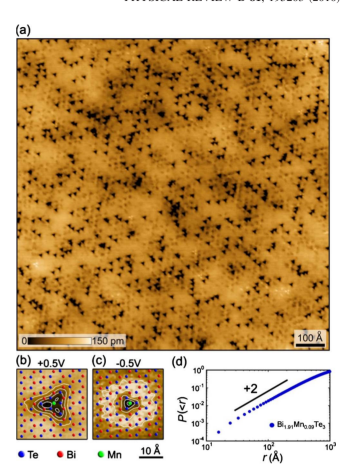


FIG. 11. (Color online) STM topographic image of Bi_{1.91}Mn_{0.09}Te₃, identifying the atomic substitutional site of Mn dopants, and the absences of Mn clustering. (a) STM topograph (+250 meV, 40 pA) of the Bi_{1.91}Mn_{0.09}Te₃ (001) surface over a 1000×1000 Å² area. Substitutional Mn atoms appear as triangular suppressions of the LDOS. [(b) and (c)] Zoom-in topographies over Mn dopants in an area of $30 \times 30 \text{ Å}^2$ for unoccupied (+500 mV, 30 pA) and filled states (-500 mV, 30 pA). The positions of top Te layer, Bi layer, and substitutional Mn are shown by blue, magenta, and green circles on the topographies. (d) Study of the possibility of clustering by calculating the probability of correlation between Mn pairs in different locations. For a randomly chosen pair of Mn, P($\langle r \rangle$ gives the probability of the pair having a separation less than r. The calculated probability scales very close to r^2 (solid line) for an extended range of distances, demonstrating the uncorrelated distribution of Mn dopants.

starting with values higher than 2 at short distances and smaller than 2 at large distances. Five topographies from randomly chosen areas of the surface several hundreds of microns apart were also analyzed and yielded similar results.

The electron acceptor nature of Mn dopants can be seen in the STM from the evolution of the LDOS with doping. The LDOS of the undoped sample, shown in Fig. 12 [blue (darker) line], consists of bulk conduction and valence band contributions that decay toward the bulk gap. Within the gap, the topological surface states of Bi_2Te_3 are the sole contributors to the LDOS. Upon Mn doping, the LDOS [red (lighter) line] shifts as a whole to higher energies, signifying the reduced density of unbound electrons in the Mn doped sample. Accordingly, the Fermi energy, E_F , shifts with doping toward

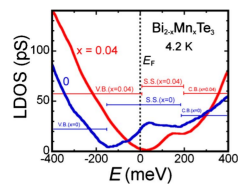


FIG. 12. (Color online) Comparison of the LDOS for pristine $\mathrm{Bi}_2\mathrm{Te}_3$ [blue (darker) line] and $\mathrm{Bi}_{1.96}\mathrm{Mn}_{0.04}\mathrm{Te}_3$ [red (lighter) line] as measured by STM at 4.2 K. Valence band (V.B.) and conduction band (C.B.) states are seen at the low and high bias values and surface states (S.S.) are seen in the band gap at intermediate bias conditions. The chemical potential is shifted by about 150 mV toward the valence band in the Mn-doped sample, indicating the p-type character of the Mn dopants. The surface states are present in Mn-doped $\mathrm{Bi}_2\mathrm{Te}_3$ below its bulk ferromagnetic T_C .

the valence band, rendering the sample *p*-type. The surface states remain present in the gap at 4.2 K in the Mn-doped crystal, well within the ferromagnetically ordered temperature regime.

Finally, we have studied the topological surface state band structure with ARPES to further characterize changes in the electronic behavior of the system on Mn doping. Previous work has shown that there is no noticeable change in the bulk valence band of Mn-doped Bi₂Te₃ when compared to pure Bi₂Te₃.9 The surface states in undoped Bi₂Te₃ are lightly electron doped, with approximately 0.001 charge carriers in the Brillioun zone, as estimated from the Fermi surface area [Figs. 13(a) and 13(b)]. The surface states remain present at the x=0.09 Mn doping level [Fig. 13(c)]. At the temperature of the measurements, 15 K, the material is very close to its full ferromagnetic transition temperature, so short-range ferromagnetic ordering is present and strong in the bulk phase. The Mn substitution results in slight hole doping of the surface states, decreasing the chemical potential by approximately 180 meV to beneath the surface state Dirac point, leaving fewer than 0.001 holelike charge carriers per surface Brillouin zone. The ARPES data show that the surface state characteristics are altered by the doping; a detailed description of doping-induced changes in the surface state dispersion will be reported elsewhere.

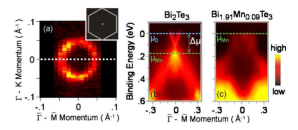


FIG. 13. (Color online) (a) Fermi surface of the surface states of undoped Bi_2Te_3 is shown, with a complete map of momentum space in the inset revealing only ~ 0.001 conducting electrons in the surface Brillouin zone. [(b) and (c)] Energy-resolved ARPES measurements show that the chemical potential of $Bi_{1.91}Mn_{0.09}Te_3$ (" μ_{Mn} ") is lowered by approximately 180 meV compared to undoped Bi_2Te_3 (" μ_0 "), resulting in some changes to the surface states and a transition from electronlike to holelike charge carriers (~ 0.001 holes/BZ) in the surface states. Data were taken approximately 30 min after cleavage.

IV. CONCLUSIONS

The development of ferromagnetism at low temperatures in Mn-doped Bi₂Te₃ has been followed as a function of dopant concentration. Ferromagnetism sets in at 2% Mn substitution for Bi, x=0.04 in Bi_{2-x}Mn_xTe₃, with a maximum T_C of about 12 K seen for $Bi_{1.91}Mn_{0.09}Te_3$. The ordered ferromagnetic moments are aligned with the easy axis perpendicular to the Bi₂Te₃ basal plane. The absence of Mn clusters indicates that the system as a true dilute ferromagnetic semiconductor. The moment per Bi_{2-x}Mn_xTe₃ formula unit is substantially larger than is observed in the analogous Fe-doped Bi₂Te₂ system.¹⁴ Important to the study of topological surface states, we have shown that for ferromagnetic samples, the surface states remain present, though altered in subtle ways. It will be of substantial future interest to characterize the impact of the magnetism on the topological surface states in this class of compounds through further measurements. The fact that doping Bi₂Se₃ with nonmagnetic Cu results in superconductivity¹² and doping Bi₂Te₃ with magnetic Mn leads to ferromagnetism opens several new doors for research on the surface states in topological insulators.

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